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Amendments to the Claims

Please amend claims 1, 8, 11, 14, 16, 18, 22-25, 32, 35, 39-41, and 44-46 as indicated in the

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listing of claims.

Please cancel claims 20 and 43 without prejudice or disclaimer.

Please add new claims 47-52.

The listing of claims will replace all prior versions, and listings of claims in the application:

Listing of Claims:

Claim 1. (Currently Amended) A method of generating a humanized mouse, comprising:

recombining a first DNA construct with a second DNA construct,

wherein the first DNA construct has a murine-mouse DNA sequence contained therein, and

wherein the second DNA construct has a human DNA sequence contained therein and the human

DNA sequence is flanked by a first and a second murine mouse DNA sequence,

wherein the first and second mouse DNA sequences are orthologous to and have the same order

and orientation relative to the human DNA sequence as human sequences flanking the human

DNA sequence when it is present in the genome of a human;

isolating a homologously recombined third DNA construct having the human DNA

sequence flanked by the first and second murine mouse DNA sequences;

introducing the recombined third DNA construct into a murine mouse embryogenic stem

cell;

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introducing the embryogenic stem cell[[s]] into a mouse blastocyst, thereby producing a chimeric blastocyst;

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implanting the chimeric blastocyst into a pseudopregnant mouse, wherein the pseudopregnant mouse delivers a humanized mouse, thereby generating a humanized mouse.

(Cancelled) Claim 2.

(Original) The method of claim 1, wherein the first DNA construct is a bacterial Claim 3. artificial chromosome.

Claim 4. (Original) The method of claim 1, wherein the second DNA construct is a bacterial artificial chromosome.

Claim 5. (Original) The method of claim 4, wherein the bacterial artificial chromosome is linearized.

Claim 6. (Original) The method of claim 1, wherein the recombining is carried out in a strain of E. coli.

(Original) The method of claim 1, wherein the E. coli is deficient for sbcB, sbcC, Claim 7. recB, recC or recD activity and has a temperature sensitive mutation in recA.

(Currently Amended) The method of claim 1, wherein the human DNA sequence Claim 8. is selected from the group consisting of a G-protein coupled receptor gene, a kinase gene, a phosphatase gene, an ion channel gene, a nuclear receptor gene, an oncogene gene, a cancer suppressor gene-gene, a viral receptor gene, a bacterial receptor gene, a P450 gene-gene, an

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insulin receptor gene, an immunoglobin gene, a metabolic pathway gene, a transcription factor gene, a hormone receptor gene, a cytokine gene, a cell signaling pathway gene and a cell cycle gene.

- (Original) The method of claim 1, wherein the third DNA construct is a bacterial Claim 9. artificial chromosome.
- Claim 10. (Original) The method of claim 1, wherein the human DNA sequence is a human gene sequence having at least one intron contained therein.
- Claim 11. (Currently Amended) The method of claim 110, wherein the third DNA construct has a selection marker contained within the at least one intron.
- Claim 12. (Original) The method of claim 11, wherein the selection marker is added following the recombining step.
- Claim 13. (Original) The method of claim 11, wherein the selection marker is a positive selection marker.
- Claim 14. (Currently Amended) The method of claim 11, wherein the third DNA construct has a second selection marker that flanks the first or the second murine mouse DNA sequence.
- Claim 15. (Cancelled)
- (Currently Amended) The method of claim 1, wherein the human DNA coding Claim 16. sequence comprises a coding sequence comprisinges a start codon having a 5' end, and the first

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mouse DNA sequence in the second construct is joined to the human DNA coding sequence at the 5' end of the start codon.

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(Previously Presented) The method of claim 16, wherein the human DNA coding Claim 17. sequence comprises a stop codon having a 3' end, and the second mouse DNA sequence in the second construct is joined to the human DNA coding sequence at the 3' end of the stop codon.

(Currently Amended) A linearized bacterial artificial chromosome DNA Claim 18. construct for performing homologous recombination within a cell of a non-human animal, the construct comprising:

a human DNA coding sequence having at least one intron disposed therein;

a selection marker gene contained within said at least one intron;

a first non-human animal DNA sequence and a second non-human animal DNA sequence, wherein said first and second non-human animal DNA sequences flank the human DNA coding sequence,

wherein the first and second non-human animal DNA sequences are from the same species, wherein the first and second non-human animal DNA sequences are orthologous to and have the same order and orientation relative to the human DNA sequence as human sequences flanking the human DNA sequence when it is present in the genome of a humanare homologous to sequences in the genome of the non-human animal that flank a gene orthologous to the human DNA coding sequence.

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(Original) The DNA construct of claim 18, further comprising a second selection Claim 19. marker adjacent to one of the non-human DNA sequences.

Claim 20. (Cancel)

Claim 21. (Original) The DNA construct of claim 18, wherein the first and second nonhuman DNA sequences are mouse genomic DNA sequences.

(Currently Amended) The DNA construct of claim 18, wherein the flanking non-Claim 22. human animal DNA sequences are from about 0.1 to 200 kb in length.

(Currently Amended) The DNA construct of claim 18, wherein the human DNA Claim 23. comprises a coding sequence comprisinges a start codon having a 5'end and the first non-human sequence is joined adjacent to the 5' end of the start codon of the human DNA coding sequence.

Claim 24. (Currently Amended) The DNA construct of claim 1823, wherein the human DNA coding sequence comprises comprises a stop codon having a 3' end and the first nonhuman sequence is joined adjacent to the 3' end of the stop codon of the human DNA coding sequence.

Claim 25. (Currently Amended). A method for generating a DNA construct for performing homologous recombination within a cell by

recombining a first DNA construct with a second DNA construct, wherein the first DNA construct has a non-human animal DNA sequence contained therein,

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wherein the second DNA construct has a human DNA sequence contained therein and the human DNA sequence is flanked by a first and a second non-human animal DNA sequence, wherein the non-human animal DNA sequences of the first and second construct are from the same species,

wherein the first and second non-human animal DNA sequences are orthologous to and have the same order and orientation relative to the human DNA sequence as human sequences flanking the human DNA sequence when it is present in the genome of a human; and

isolating a homologously recombined third DNA construct having the human DNA sequence flanked by the first and second non-human animal DNA sequence.

Claim 26. (Cancelled)

Claim 27. (Original) The method of claim 25, wherein the first DNA construct is a bacterial artificial chromosome.

Claim 28. (Original) The method of claim 25, wherein the second DNA construct is a bacterial artificial chromosome.

(Original) The method of claim 28, wherein the bacterial artificial chromosome is Claim 29. linearized.

(Original) The method of claim 25, wherein the recombining is carried out in a Claim 30. strain of E. coli.

Claim 31. (Original) The method of claim 25, wherein the E. coli is deficient for sbcB, sbcC, recB, recC or recD activity and has a temperature sensitive mutation in recA.

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Claim 32. (Currently Amended) The method of claim 25, wherein the human DNA

sequence is selected from the group consisting of a G-protein coupled receptor gene, a kinase

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gene, a phosphatase gene, an ion channel gene, a nuclear receptor gene, an oncogene-gene, a

cancer suppressor gene-gene, a viral receptor-gene, a bacterial receptor gene, a P450 gene gene,

an insulin receptor gene, an immunoglobin gene, a metabolic pathway gene, a transcription

factor gene, a hormone receptor gene, a cytokine gene, a cell signaling pathway gene and a cell

cycle gene.

(Original) The method of claim 25, wherein the third DNA construct is a bacterial Claim 33.

artificial chromosome.

Claim 34. (Original) The method of claim 25, wherein the human DNA sequence is a human

gene sequence having at least one intron contained therein.

Claim 35. (Currently Amended) The method of claim 2534, wherein the third DNA

construct has a selection marker contained within the at least one intron.

(Original) The method of claim 35, wherein the selection marker is added Claim 36.

following the recombining step.

(Original) The method of claim 35, wherein the selection marker is a positive Claim 37.

selection marker.

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(Previously Presented) The method of claim 35, wherein the third DNA construct Claim 38.

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has a second selection marker that flanks the first or the second non-human animal DNA

sequence.

Claim 39. (Currently Amended) The method of claim 25, wherein the human DNA

sequence comprises a coding sequence comprising a start codon having a 5' end, and the first

non-human DNA sequence in the second construct is joined to the human DNA coding sequence

at the 5' end of the start codon.

(Currently Amended) The method of claim 39, wherein the human DNA coding Claim 40.

sequence comprises a stop codon having a 3' end, and the second non-human DNA sequence in

the second construct is joined to the 3' of the stop codon.

Claim 41. (Currently Amended) A humanized mouse produced by the method of claim 1,

wherein the human DNA sequence is a gene selected from a G-protein coupled receptor gene, a

kinase gene, a phosphatase gene, an ion channel gene, a nuclear receptor gene, an oncogene

gene, a cancer suppressor gene-gene, a viral receptor gene, a bacterial receptor gene, a P450 gene

gene, an insulin receptor gene, an immunoglobin gene, a metabolic pathway gene, a

transcription factor gene, a hormone receptor gene, a cytokine gene, a cell signaling pathway

gene and a cell cycle gene.

Claim 42. (Cancelled)

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Claim 43. (Previously Presented) The humanized mouse of claim 41, wherein the gene is

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involved in human drug metabolism.

Claim 44. (Currently Amended) The humanized mouse of claim 41, wherein the gene is a

PXR, CAR, RXR, CYP3A4, CYP2B6, CYP2C9 or MDR1 gene.

(Currently Amended) A method of generating a humanized cell, comprising: Claim 45.

recombining a first DNA construct with a second DNA construct,

wherein the first DNA construct has a non-human animal DNA sequence contained therein, and

wherein the second DNA construct has a human DNA sequence contained therein and the human

DNA sequence is flanked by a first and a second non-human animal DNA sequence,

wherein the first and second non-human animal DNA sequences are from the same species,

wherein the first and second non-human animal DNA sequences are orthologous to and have the

same order and orientation relative to the human DNA sequence as human sequences flanking

the human DNA sequence when it is present in the genome of a human;

isolating a homologously recombined third DNA construct having a the human DNA

sequence flanked by the first and second non-human animal DNA sequence; and

introducing the recombined third DNA construct into a non-human animal cell of the

same species as the non-human DNA sequences of the first and second constructs, thereby

generating a humanized cell.

(Currently Amended) The method of claim 45, wherein the non-human animal Claim 46.

cell is an mouse embryogenic stem cell.

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(New) A DNA construct for performing homologous recombination within a cell Claim 47.

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of a non-human animal, the construct comprising:

a human DNA coding sequence having at least one intron disposed therein;

a selection marker gene contained within said at least one intron;

a first non-human animal DNA sequence and a second non-human animal DNA

sequence,

wherein said first and second non-human animal DNA sequences flank the human DNA coding

sequence,

wherein the first and second non-human animal DNA sequences are from the same species,

wherein the first and second non-human animal DNA sequences are orthologous to and have the

same order and orientation relative to the human DNA sequence as human sequences flanking

the human DNA sequence when it is present in the genome of a human; and

a second selection marker adjacent to one of the non-human DNA sequences.

(New) The DNA construct of claim 47, wherein the construct is a linearized Claim 48.

bacterial artificial chromosome.

(New) The DNA construct of claim 47, wherein the first and second non-human

DNA sequences are mouse genomic DNA sequences.

Claim 50. (New) The DNA construct of claim 47, wherein the non-human animal DNA

sequences are from about 0.1 to 200 kb in length.

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Claim 51. (New) The DNA construct of claim 47, wherein the human DNA sequence comprises a coding sequence comprising a start codon having a 5'end and the first non-human sequence is joined adjacent to the 5' end of the start codon of the human DNA coding sequence.

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Claim 52. (New) The DNA construct of claim 51, wherein the human DNA coding sequence comprises a stop codon having a 3' end and the first non-human sequence is joined adjacent to the 3' end of the stop codon of the human DNA coding sequence.